Proteomic analysis of the *S. cerevisiae* response to the anticancer ruthenium complex KP1019

Laura K. Stultz  
*Birmingham-Southern College*

Alexandra Hunsucker  
*Birmingham-Southern College*

Sydney Middleton  
*Birmingham-Southern College*

Evan Grovenstein  
*Birmingham-Southern College*

Jacob O'Leary  
*Birmingham-Southern College*

*See next page for additional authors*

Follow this and additional works at: https://scholarexchange.furman.edu/oa-fund

Part of the Medicine and Health Sciences Commons

**Recommended Citation**  

This Article (Journal or Newsletter) is made available online by part of the Furman University Scholar Exchange (FUSE). It has been accepted for inclusion in Open Access Fund Publications by an authorized FUSE administrator. For terms of use, please refer to the FUSE Institutional Repository Guidelines. For more information, please contact scholarexchange@furman.edu.
Proteomic analysis of the *S. cerevisiae* response to the anticancer ruthenium complex KP1019

Laura K. Stultz,*a* Alexandra Hunsucker, b Sydney Middleton, a Evan Grovenstein, b Jacob O’Leary, a Eliot Blatt, c Mary Miller, c James Mobley d and Pamela K. Hanson e

Like platinum-based chemotherapeutics, the anticancer ruthenium complex indazolium trans-[tetra-chlorobis(1H-indazole)ruthenate(III)], or KP1019, damages DNA, induces apoptosis, and causes tumor regression in animal models. Unlike platinum-based drugs, KP1019 showed no dose-limiting toxicity in a phase I clinical trial. Despite these advances, the mechanism(s) and target(s) of KP1019 remain unclear. For example, the drug may damage DNA directly or by causing oxidative stress. Likewise, KP1019 binds cytosolic proteins, suggesting DNA is not the sole target. Here we use the budding yeast *Saccharomyces cerevisiae* as a model in a proteomic study of the cellular response to KP1019. Mapping protein level changes onto metabolic pathways revealed patterns consistent with elevated synthesis and/or cycling of the antioxidant glutathione, suggesting KP1019 induces oxidative stress. This result was supported by increased fluorescence of the redox-sensitive dye DCFH-DA and increased KP1019 sensitivity of yeast lacking Yap1, a master regulator of the oxidative stress response. In addition to oxidative and DNA stress, bioinformatic analysis revealed drug-dependent increases in proteins involved ribosome biogenesis, translation, and protein (re)folding. Consistent with proteotoxic effects, KP1019 increased expression of a heat-shock element (HSE) *lacZ* reporter. KP1019 pre-treatment also sensitized yeast to oxaliplatin, paralleling prior research showing that cancer cell lines with elevated levels of translation machinery are hypersensitive to oxaliplatin. Combined, these data suggest that one of KP1019’s many targets may be protein metabolism, which opens up intriguing possibilities for combination therapy.

Significance to metallomics

The ruthenium complex KP1019 has promising anticancer properties, yet its mechanisms of action are only partially understood. Using budding yeast as a model system, here we report the first global proteomic analysis examining the cellular response to this drug. One of the strongest signatures revealed by this work was elevated levels of proteins involved in ribosomal biogenesis (rbi) and translation, suggesting that KP1019 perturbs protein homeostasis. Previous studies showed that enhanced rbi gene expression is associated with increased sensitivity to oxaliplatin. Our observation of synergy between KP1019 and oxaliplatin suggests future promise for combination therapy.

Introduction

Certain metals, including iron and copper, serve essential functions in cells. However, at high levels, many metals are toxic.

Although mechanisms of toxicity can vary, leading to divergent cellular responses,1,2 many metals have been shown to induce oxidative stress and DNA damage.3–6 Some metals have also been shown to induce protein misfolding and aggregation.7

In recent decades, the toxicity of a diverse array of metals has been harnessed in the production of anti-cancer drugs.8–11 Early examples include cisplatin and carboplatin, which are often used to treat testicular and ovarian cancers, as well as head and neck tumours. More recently, oxaliplatin has emerged as an effective treatment for gastrointestinal cancers,12,13 and it may be particularly effective in the context of combination therapies.14 Unfortunately, the utility of platinum-based drugs can be hindered by dose-limiting toxicity and development of drug resistance.15–17

---

* a Department of Chemistry, Birmingham-Southern College, Birmingham, AL 35254, USA
b Department of Biology, Birmingham-Southern College, Birmingham, AL 35254, USA
c Department of Biology, Rhodes College, Memphis, TN 38112, USA
d Department of Surgery, University of Alabama at Birmingham, School of Medicine, Birmingham, AL 35294, USA
e Department of Biology, Furman University, Greenville, SC 29613, USA.
E-mail: pamela.hanson@furman.edu

Received 8th January 2020, Accepted 18th March 2020
DOI: 10.1039/d0mt00008f
rsc.li/metallomics
Though not yet approved for regular use, three ruthenium complexes – NAMI-A, KP1019, and KP1339 – show significant promise in preclinical studies and early clinical trials.\textsuperscript{18,19} For example, indazolium \textit{trans-[tetrachlorobis(1H-indazole)ruthenate(III)]}, or KP1019 (Fig. 1), displays no dose-limiting toxicity\textsuperscript{20,21} and maintains potency against drug resistant cell lines.\textsuperscript{22} KP1019 caused tumour regression in animal models,\textsuperscript{23} and in a phase I clinical trial, the drug stabilized disease progression in five of six evaluable patients.\textsuperscript{20,21} However, this drug’s mechanism of action remains poorly understood. In colorectal cancer cells, KP1019 has been shown to induce oxidative stress and DNA damage,\textsuperscript{24} yet in cervical carcinoma cells, the drug bound primarily to proteins in the cytosolic fraction.\textsuperscript{25} Furthermore, KP1019 not only bound genomic DNA but also accumulated in mitochondria of ovarian cancer cells.\textsuperscript{26}

Consistent with findings in cancer cell lines, KP1019 has been shown to inhibit growth, induce cell death, and damage DNA in the budding yeast \textit{Saccharomyces cerevisiae}.\textsuperscript{27–29} Moreover, the drug bound readily with cytoplasmic proteins and mitochondria of fractionated yeast cells.\textsuperscript{30} KP1019’s ability to induce stress is also consistent with its ability to activate the evolutionarily conserved stress-responsive kinase Hog1\textsuperscript{31} and to increase lipid droplet formation.\textsuperscript{22} Combined, these studies suggest that KP1019 has multiple modes of action in yeast, much like it has in cultured mammalian cells. Thus, budding yeast appears to be an appropriate model organism for better characterizing KP1019’s impact on cells.

When working with poorly characterized drugs, unbiased genome-wide approaches can help establish the targets of bioactive compounds.\textsuperscript{33,34} For example, transcriptional profiling of yeast helped uncover quinine’s ability to perturb glucose transport\textsuperscript{35} and contributed to establishment of calcium mobility as a mechanism of action for the antifungal terpenoid phenol carvacrol.\textsuperscript{36}

Given the success of such global approaches, we have used mass spectrometry to analyse the effects of KP1019 on the budding yeast proteome. Our results support previous findings, verifying activation of the DNA damage response, while revealing new physiological responses to the drug, such as morphogenesis, metabolic re-tooling, and induction of ribosomal biogenesis.

### Materials and methods

#### Yeast strains and growth conditions

Yeast strains used in this study are listed in Table 1. Yeast were grown under standard conditions, at 30 °C using the rich medium yeast extract peptone dextrose (YPD) (1% yeast extract, 2% bacto-peptone, 2% dextrose) or synthetic complete media (SDC) as indicated.\textsuperscript{39} For experiments involving plasmids, a standard lithium acetate transformation protocol\textsuperscript{40} was used, and transformants were selected on minimal media lacking the component necessary for plasmid maintenance.

#### Drug synthesis

KP1019 was synthesized using a protocol described previously\textsuperscript{27} and adapted from Lipponer \textit{et al.}.\textsuperscript{41} Briefly, 1 g RuCl$_3$ $\cdot$ 3H$_2$O was added to 20 ml of 12 M HCl and 20 ml of ethanol and refluxed for 1 hour. Once the reaction mixture had cooled, the ethanol was removed using a rotary evaporator, and 12 M HCl was added to give a final volume of 40 ml. The ruthenium solution was combined with 3.74 g of indazole that had been dissolved in 60 ml of 12 M HCl at 70 °C. This reaction mixture was heated for 15 minutes at 80–90 °C and cooled to room temperature with stirring. The resulting solid was filtered, and then stirred for 2 hours at room temperature for 2 hours. The solid was filtered and subsequently washed with cold ethanol followed by cold diethyl ether. The drug was dried under vacuum for 18–24 hours. KP1019 purity was verified by UV-visible spectroscopy, elemental analysis and cyclic voltammetry.

#### Proteomic analysis

Proteomic analysis was carried out as previously,\textsuperscript{42} with minor changes included below.  

**Sample preparation.** For each of three biological replicates, 250 ml of wild-type yeast (BY 4742) were grown to mid-log phase
Metallomics, 2020, 12, 876–890

(OD_{600} 0.5–1.0) at 30 °C in SDC. The culture was then split and half was treated with 80 µg ml^{-1} KP1019 for 3 hours; the other half was left untreated. Samples were washed with water and cell concentrations were normalized. Cells were then pelleted and stored at −80 °C. The Y-PER Yeast Protein Extraction Reagent (Thermo Fisher, Waltham, MA) was used to extract proteins from the pellets. The protein fractions were quantified using Pierce BCA Protein Assay Kit (Thermo Fisher Scientific, Cat.# PI23223), ~20 µg of protein per sample were then diluted to 35 µl using NuPAGE LDS sample buffer (1× final conc., Invitrogen, Cat.#NP0007). Proteins were then reduced with DTT and denatured at 70 °C for 10 min prior to loading everything onto Novex NuPAGE 10% Bis-Tris protein gels (Invitrogen, Cat.# NP0315BOX) and separated as a short stack (10 min at 200 constant V). The gels were stained overnight with Novex Colloidal Blue Staining kit (Invitrogen, Cat.# LC6025).

Following de-staining, each lane was cut into single MW fractions and equilibrated in 100 mM ammonium bicarbonate (AmBc), each gel plug was then digested overnight with Trypsin Gold, Mass Spectrometry Grade (Promega, Cat.# V5280) following manufacturer’s instruction. Peptide extracts were reconstituted in 0.1% formic acid/ddH_{2}O at 0.1 µg µl^{-1}.

**Mass spectrometry.** Peptide digests (8 µL each) were injected onto a 1260 Infinity nHPLC stack (Agilent Technologies), and separated using a 75 micron I.D. × 15 cm pulled tip C-18 column (Jupiter C-18 300 Å, 5 micron, Phenomenex). This system runs in-line with a Thermo Orbitrap Velos Pro hybrid mass spectrometer, equipped with a nano-electrospray source (Thermo Fisher Scientific), and all data were collected in CID mode. The nHPLC was configured with binary mobile phases that included solvent A (0.1% FA in ddH_{2}O), and solvent B (0.1% FA in 15% ddH_{2}O/85% ACN), programmed as follows; 10 min @ 5%B (2 µl min^{-1}, load), 90 min @ 5–40% B (linear: 0.5 nl min^{-1}, analyse), 5 min @ 70% B (2 µl min^{-1}, wash), 10 min @ 0% B (2 µl min^{-1}, equilibrate). Following each parent ion scan (300–1200 m/z @ 60k resolution), fragmentation data (MS2) was collected on the top most intense 15 ions. For data dependent scans, charge state screening and dynamic exclusion were enabled with a repeat count of 2, repeat duration of 30 s, and exclusion duration of 90 s.

**MS data conversion and searches.** The XCalibur RAW files were collected in profile mode, centroided and converted to MzXML using ReAdW v. 3.5.1. The mgf files were then be created using MzXML2Search (included in TPP v. 3.5) for all scans. The data was searched using SEQUEST, which was set for two maximum missed cleavages, a precursor mass window of 20 ppm, trypsin digestion, variable modification C @ 57.0293, and M @ 15.9949. Searches were performed with a species specific subset of the UniRef100 database.

**Peptide filtering, grouping, and quantification.** The list of peptide IDs generated based on SEQUEST (Thermo Fisher Scientific) search results were filtered using Scaffold (Protein Sciences, Portland Oregon). Scaffold filters and groups all peptides to generate and retain only high confidence IDs while also generating normalized spectral counts (N-SC’s) across all samples for the purpose of relative quantification. The filter cut-off values were set with minimum peptide length of > 5 AA’s, with no MH + 1 charge states, with peptide probabilities of > 80% C.I., and with the number of peptides per protein ≥ 2. The protein probabilities were then set to a > 99.0% C.I., and an FDR < 1.0. Scaffold incorporates the two most common methods for statistical validation of large proteome datasets, the false discovery rate (FDR) and protein probability.43–45 Relative quantification across experiments were then performed via spectral counting,46,47 and when relevant, spectral count abundances were then normalized between samples.48

**Systems analysis.** Gene ontology assignments, clustering, and pathway analysis were carried out using Search Tool for Recurring Instances of Neighbouring Genes (STRING; https://string-db.org/),49 Protein Analysis TThrough Evolutionary Relationships (PANTHER; http://www.pantherdb.org/),50 and YeastMine (http://yeastmine.yeastgenome.org).51

**Filamentation assay.** A wild-type strain from the filamentation competent Σ1278b background52,53 was cultured overnight in SDC then spread onto SDC media containing the indicated concentrations of KP1019. After an 18 hour incubation at 30 °C colony morphology was documented using a Zeiss Axioskop 2 plus with Zeiss Axiovision software (Carl Zeiss AG, Oberkochen, Germany).

**Oxidative stress assay.** Wild-type yeast were cultured to mid-log phase then incubated with 50 µM 2,7’-dichlorodihydrofluorescein diacetate (DCFH-DA; Sigma-Aldrich, St. Louis, MO) for 1.5 hours at 30 °C to pre-load the cells with the redox sensitive dye. The samples were then treated with 0 or 80 µg ml^{-1} KP1019 or 10 mM H_{2}O_{2} for 2.5 hours. Samples were then analysed by flow cytometry using a BD Accuri C6 Flow Cytometer (Becton Dickson, Franklin Lakes, NJ). 10 000 events were collected for each sample.

**Beta-galactosidase assays.** The genotoxicity and proteotoxicity of KP1019 were verified using wild-type yeast strain BY4742 transformed with the pZ22 RNR3-lacZ or HSE-lacZ reporter constructs, respectively.54,55 Specifically, transformed cells were cultured to mid-log phase (OD_{600} 0.5–1.0) in selective media then treated with KP1019 or 10 mM H_{2}O_{2} dissolved in the same type of medium. Samples were incubated for 3 hours to allow for gene induction, at which point β-galactosidase activity was measured using the permeabilized cell assay described by Guarante.56

**Heat shock assay.** The heat shock sensitivity assay was adapted from Wang et al.57 Briefly, yeast were cultured to mid-log phase at 30 °C in SDC then treated with the indicated concentration of KP1019 for 2 hours. Cells were then washed with SDC and cell concentrations were normalized. Samples were then incubated at 32 °C, and aliquots were removed and pipetted onto YPD at the indicated times. Following a 48 hour incubation at 30 °C, images were captured using a scanner.
Drug sensitivity and synergy assays

Drug-containing SDC was subjected to 2-fold serial dilution across the wells of a microtitre plate. Overnight cultures of yeast were diluted to OD\textsubscript{600} 0.1 and subsequently diluted 20-fold more in SDC. An equal volume of diluted cell suspension was added to each well of the microtitre plate. After an 18 to 24 hour incubation at 30 °C, the growth of each strain at each concentration of drug was recorded as absorbance at 630 nm using a BioTek (Winooski, VT) microtitre plate reader. Data were fit with a sigmoidal 4-parameter logistic curve to determine the concentration of drug that inhibited growth by 50%.

For the synergy assay, overnight cultures of yeast were subcultured to mid-log phase (OD\textsubscript{600} 0.5–1.0) in SDC prior to transferring the cells to SDC containing the indicated concentrations of KP1019. Following a three-hour incubation with the drug, cells were harvested by centrifugation then resuspended in drug-free SDC. Samples were then normalized to OD\textsubscript{600} 0.1 and subsequently diluted 10-fold more in SDC. One part yeast suspension was then added to three parts oxaliplatin in SDC to achieve the desired final concentration of platinum drug. After an 18 to 24 hour incubation at 30 °C, yeast growth at each concentration of drug was recorded as described above. The resulting data were analysed using SynergyFinder.\textsuperscript{58}

Results

Analysis of KP1019-dependent changes in protein abundance

To characterize the cellular response to the anticancer ruthenium complex KP1019, we used a proteomic approach to measure changes in protein abundance in yeast treated with 80 μg ml\textsuperscript{-1} KP1019, a concentration used in prior transcriptomic study\textsuperscript{29} and previously shown to robustly inhibit yeast growth and arrest the cell cycle while causing only modest amounts of cell death.\textsuperscript{27} The yeast proteome was analysed after incubating the cells with KP1019 for 3 hours,\textsuperscript{29} the same duration of exposure used in prior transcriptomic studies of this drug.\textsuperscript{29,32} Ultimately, 3676 different proteins were detected across three biological replicates. To determine which proteins displayed a significant change in expression level, a student’s t-test was performed and proteins with p-values less than 0.05 were considered to display statistically significant differences in abundance. In an attempt to enrich for physiologically significant changes in protein levels, we limited further analysis to proteins with a 1.5-fold or greater increase in abundance or a 25% or greater decrease in abundance, consistent with a prior study of stress-dependent changes in protein levels in yeast.\textsuperscript{60} Using these criteria, levels of 230 proteins increased and 200 proteins decreased following a three-hour incubation with the drug (Fig. 2).

Amongst these 430 proteins, fold change ranged from a high of 24-fold repression for the spore wall protein Irc18.\textsuperscript{61} To identify transcription factors that may contribute to KP1019-dependent changes in protein levels, the pool of induced and repressed proteins were analysed in Yeastract (http://www.yeastract.com).\textsuperscript{63} Specifically, we applied Yeastract’s TFRank functionality with a heat diffusion coefficient of 0.25 to avoid focusing on overly proximal or distal layers of regulation.\textsuperscript{64} Implicated transcription factors are presented in Fig. 5A. Since four of the top ten weighted transcription factors were induced and reduced abundance proteins both had significantly more interactions than expected with protein–protein interaction (PPI) enrichment p-values of 2 × 10\textsuperscript{-8} and 4 × 10\textsuperscript{-7}, respectively. However, these proteins were included in subsequent bioinformatics analyses but are not included in the volcano plot.
are known for their roles in filamentous growth and/or morphogenesis, we conducted a preliminary validation of the TFRank analysis by plating the filamentation competent strain \( \Sigma 1278b^{52,53} \) on media containing varying concentrations of KP1019. As seen in Fig. 5B, the drug induced a dose-dependent increase in morphogenesis.

As will be discussed in more detail below, many of the findings generated through bioinformatics analyses revealed
consistent themes, including activation of multiple stress response pathways. In fact, one of the largest clusters identified by STRING contains proteins involved in diverse biological processes including cellular detoxification (e.g. – Sfa1), RNA metabolism (e.g. – Rtc3), and signal transduction (e.g. – Tfs1) (Fig. 3A). Though seemingly dissimilar, most have been shown to be stress-responsive, increasing in abundance and/or changing localization following application of environmental stressors. For example, the glycogen branching enzyme Glc3, was previously shown to increase in abundance and form cytoplasmic puncta upon induction of DNA replication stress.

KP1019 activates oxidative stress responses

Given that KP1019 has previously been shown to induce oxidative stress in colorectal carcinoma cells, it was not surprising to find “oxidation–reduction process” and “glutathione metabolism” amongst the enriched GO terms and Kyoto Encyclopedia of Genes and Genomes (KEGG) pathways, respectively (Fig. 3). Consistent with these findings, bioinformatics analyses uncovered additional evidence of metabolic retooling. For example, when proteins displaying drug-dependent reductions in abundance were analysed by STRING, “metabolic pathways” was identified as a significantly enriched KEGG pathway (FDR = 0.0001). Likewise, PANTHER revealed enrichment of proteins involved in purine nucleotide and alpha amino acid metabolism (Fig. 4B).

To better integrate these bioinformatics findings, metabolic enzymes displaying drug-dependent changes in abundance were mapped onto several intertwined metabolic pathways (Fig. 6A). This visualization reveals how these metabolic changes may coordinately regulate increased synthesis and/or cycling of the cellular antioxidant glutathione. In addition to increasing abundance of enzymes (Gsh2, Gtt1, and Glr1) directly involved in glutathione synthesis/cycling, KP1019 treatment increased levels of several enzymes in the pentose phosphate pathway (PPP), including the glucose-6-phosphate dehydrogenase Zwf1, which catalyses the pathway’s rate-limiting step. Moreover, drug exposure lowered levels of proteins involved in inositol synthesis, potentially reducing the shunting of glucose-6-phosphate out of the PPP. Increased flux through the PPP could increase levels of NADPH, which is critical for glutathione regeneration. Furthermore, YeastMine uncovered significant changes to the S-adenosylmethionine (SAM) cycle (Benjamini–Hochberg corrected p-value = 0.002), which can be visualized as a cluster in Fig. 4A. This drug-dependent down-regulation might also contribute to antioxidant synthesis by increasing the proportion of de novo homocysteine available for conversion to cysteine and incorporation into GSH.

To more directly measure KP1019’s ability to induce oxidative stress, we used the redox sensitive fluorophore DCFH-DA. As seen in Fig. 6B, KP1019 treated cells displayed elevated levels of fluorescence, consistent with drug-induced oxidative stress, albeit significantly weaker than the hydrogen peroxide positive control.

To gather independent confirmation of oxidative stress, we leveraged the results of the TFRank analysis, which implicated Yap1 (Fig. 5A), a major regulator of the yeast oxidative stress response and the yeast orthologue of the human transcription factor AP-1. Specifically, we examined the KP1019 sensitivity of yeast lacking YAP1. As seen in Fig. 6C, we confirmed that yap1Δ yeast are extremely sensitive to H2O2. Loss of YAPI also caused a modest but statistically significant increase in sensitivity to KP1019, a result consistent with the modest induction of DCFH-DA fluorescence noted previously.

KP1019 induces the DNA damage response

Since several studies have shown that KP1019 damages DNA, it was surprising that the DNA damage response (DDR) was not among the enriched biological process GO terms. However, the
The ribonucleoside-diphosphate reductase (RNR) complex is visible in the STRING clustering (Fig. 3A), and it was one of the few specific cellular component GO terms that was significantly enriched (FDR = 0.01) for the list of proteins with increased abundance following KP1019 treatment. The RNR genes are well established as downstream effectors in the yeast DDR.\textsuperscript{78,79} To verify that elevated RNR levels in KP1019 treated cells was due to activation of the DDR, we examined expression of an RNR3-lacZ reporter construct in yeast lacking \textit{DDC1} and \textit{DUN1}, which encode key mediators of the DDR.\textsuperscript{80–82} As can be seen in Fig. 7A, KP1019-dependent induction of this reporter for genotoxicity\textsuperscript{83} was blocked by disruption of the DDR. Consistent with previous studies,\textsuperscript{28,29} activation of the DDR appears to be physiologically significant, as deletion of \textit{DDC1} and \textit{DUN1} increases sensitivity to KP1019, decreasing the concentration of drug that inhibits growth by 50\% (IC\textsubscript{50}) from 6.8 (wild-type) to 3.6 and 3.0 \textit{mM} for \textit{dun1\textsuperscript{D}} and \textit{ddc1\textsuperscript{D}}, respectively (Fig. 7B).

**KP1019 affects protein synthesis and folding**

Among the most consistent themes that emerged from analysis of the proteomic data were protein (re)folding and protein synthesis. As can be seen in Fig. 3, KP1019 elevated the levels of chaperones that assist with protein (re)folding. Interestingly, subjecting the induced protein dataset to publication enrichment analysis in YeastMine revealed a potential connection to the heat shock response. Specifically, three of the top 5 hits (Benjamini Hochberg corrected \textit{p}-values < 0.0001) were published articles focused on the heat shock response. In yeast, the heat shock response (HSR) involves elevated expression of chaperones. Consistent with activation of the HSR, we observed KP1019-dependent increases in the abundance of several heat shock proteins, including the HSP70 family members Ssa1-4 and the yeast HSP90 chaperones Hsp82 and Hsc82 (Fig. 3A). To determine whether the HSR transcriptional response might account for the changes in chaperone levels observed in our data, we monitored expression of a lacZ reporter under control of a heat shock element (HSE).\textsuperscript{84} Consistent with activation of the HSR, we observed a small but statistically significant drug-dependent increase in HSE-lacZ expression (Fig. 8A). Although a dose–response curve indicated that 80 \textit{\mu M} KP1019 (134 \textit{\mu M}) results in maximal activation of this reporter construct (data not shown), the ability of 80 \textit{\mu M} KP1019 to induce expression was significantly lower than that of 100 \textit{\mu M} cadmium (Fig. 8A).
This concentration of cadmium was previously shown to derepress Hsf1, the evolutionarily conserved transcription factor and master regulator of \textit{S. cerevisiae} HSR.\textsuperscript{75} Consistent with the modest induction of \textit{HSE-lacZ} by KP1019, Hsf1 was only weakly implicated in the TFRank analysis of KP1019 response. Specifically, this transcription factor ranked 30th amongst the 100+ potential regulators reported by Yeastract. To determine whether the KP1019-dependent induction of chaperones was likely to be physiologically significant, we pre-treated yeast with KP1019 and then tested the yeast for heat tolerance. As can be seen in Fig. 8B, pre-treatment with KP1019 increased yeast tolerance to heat shock.

In addition to increasing abundance of proteins involved in protein (re)folding, clusters of ribosomal and ribosomal biogenesis proteins exhibited elevated levels (Fig. 3A). Specifically, the abundance of ribosomal and ribosomal biogenesis proteins in these intertwined clusters were increased an average of four-fold (range = 1.6 to 13-fold, not including Utp9, which was not detected in the absence of KP1019) Consistent with the visualization depicted in Fig. 3A, STRING’s analytics revealed statistically significant (FDR = 0.002) enrichment of proteins associated with the cellular component GO term “cytosolic ribosome”. Likewise Yeastract identified the transcription factor Sfp1 – a major regulator of ribosomal gene expression\textsuperscript{85} – as a potential contributor to KP1019-dependent changes in protein abundance (Fig. 5A). Notably, the levels of Sfp1 itself increased four-fold in the drug treated samples.

Since elevated levels of translation machinery correlate with increased oxaliplatin sensitivity,\textsuperscript{86} and since KP1019 increases abundance of several ribosomal proteins (Fig. 3), we hypothesized that KP1019 might increase oxaliplatin sensitivity. As seen in Fig. 9A, treating yeast with 80 \(\mu\text{g mL}^{-1}\) KP1019 for three hours resulted in a roughly 60% decrease in the IC\textsubscript{50} for oxaliplatin. To determine whether this effect was synergistic, we examined a wider range of KP1019 concentrations (Fig. 9B), analysing the resulting growth inhibition with the zero interaction potency (ZIP) model, which has a relatively low false positive rate and avoids the assumptions associated with more traditional additivity and independence models.\textsuperscript{87} According to this model a delta (\(\delta\)) score of zero suggests no interaction between two drugs, whereas scores greater than zero indicate synergy and scores below zero correspond to antagonism. Although the average \(\delta\) of 5.18 suggests only modest synergy between KP1019 and oxaliplatin, within the larger interaction landscape there is a region of synergy at moderately high doses of both drugs, peaking at \(\delta = 25\) (Fig. 9C).

**Discussion**

In this study, we used the budding yeast \textit{S. cerevisiae} as a model organism to examine the global effects of the promising anticancer ruthenium complex KP1019 at 80 \(\mu\text{g mL}^{-1}\), a concentration used in a prior transcriptomic study in yeast.\textsuperscript{29} Notably, this concentration is less than two-fold higher than the maximum plasma concentration (51.5 \(\mu\text{g mL}^{-1}\)) observed in a Phase I clinical trial.\textsuperscript{20} Using a slightly higher concentration in \textit{S. cerevisiae} is appropriate, as yeast often have elevated levels of resistance to antineoplastic agents.\textsuperscript{88}
Consistent with previous findings, our results demonstrated that the drug significantly increased levels of proteins involved in genotoxic, proteotoxic, and oxidative stress responses.

With respect to the drug's impact on DNA, KP1019 binds model nucleotides in vitro and damages DNA in cancer and yeast cells; in S. cerevisiae, the resulting activation of the DNA damage response (DDR) leads to cell cycle arrest. Here, we observed induction of the ribonucleotide reductase complex (Fig. 3A) in a DDR-dependent manner (Fig. 7), and there was no evidence to suggest that these or other differences in protein levels were due solely to the drug's impact on cell cycle progression. These new data supporting KP1019-dependent activation of the DDR confirm prior studies, while validating our experimental design. We also noted that the drug induced filamentous growth in yeast (Fig. 5B), which may be a sign of replication stress.

Interestingly, DNA damage and repair were not among the enriched GO terms identified in our bioinformatics analyses. Some lack of congruence with recent transcriptomic studies would be expected due to differences that arise when characterizing proteomes as opposed to transcriptomes. For example, osmotic or oxidative insults have been shown to impact translation, at times resulting in changes to the abundance of specific proteins. Divergence from transcriptomic studies may also stem from technical reasons. Although all three studies treated yeast with KP1019 for three hours prior to analysis, discrepancies included use of different wild-type yeast strains and varying concentrations of KP1019 (80 μg ml⁻¹ [here and Bierle et al. 29] vs. 50 μg ml⁻¹ [32]), and different cut-offs for bioinformatic analyses (e.g. – 1.5 [here and Golla et al. 32] vs. 3.5 fold induction). Solubilizing the drug in DMSO, as in Golla et al. 32 but not here, may also be a confounding variable, as DMSO impacts KP1019’s stability (Cetto A and Stultz LK, personal communication). Regardless of the etiology of the variation between studies, our results support the existence of drug targets besides DNA. In particular, network and GO term analyses are consistent with the drug impacting proteins and redox processes.

In support of drug-dependent oxidative stress, we observed increased abundance of proteins involved in the pentose phosphate pathway and GSH synthesis (Fig. 3 and 6). Though these results are aligned with the oxidative stress observed in mammalian studies, increased expression of oxidative stress genes was not reported in the previous transcriptomic analyses in yeast. Regardless, we were able to verify induction of oxidative stress by measuring drug-dependent increases in fluorescence of the redox-sensitive dye DCFH-DA (Fig. 6B) and modest increases in sensitivity of yeast lacking Yap1 (Fig. 6C), the master regulator of the S. cerevisiae oxidative stress response. These findings are consistent with the “activation by reduction” hypothesis, according to which, KP1019 is a pro-drug that becomes bioactive after its central ruthenium atom is reduced from Ru³⁺ to Ru²⁺. Unlike most platinum-based drugs, which do not change oxidation states, KP1019 is more likely to be reduced in the low oxygen microenvironment of a poorly vascularized solid tumour. This localized activation may contribute to this ruthenium complex’s selectivity for malignant tissues. An anticipated byproduct of KP1019 reduction would be reactive oxygen species (ROS), which could help mediate the drug’s toxic effects. Although the more dramatic sensitivity of DDR mutants (Fig. 7B) relative to the Yap1 deletion strain (Fig. 6C) suggests DNA damage plays a greater role in the drug’s toxicity, the physiological significance and origins of drug-induced ROS require further investigation, as the oxidative stress and DNA damage responses are highly intertwined. For example, oxidative stress is a well-known cause of DNA damage, and DNA damage leads to signalling-related increases in ROS.

In addition to verifying KP1019’s ability to induce DNA damage and oxidative stress, here we showed significant increases in levels of proteins involved in protein folding (Fig. 3). In fact, the chaperone Hsp104 showed the greatest fold increase in abundance following...
drug treatment (Fig. 2). Although analogous findings were not reported in previous transcriptomic studies,25,32 these observations are consistent with KP1019’s reported binding to proteins in the cytosolic fraction of cancer cells.25 The drug’s effects on proteins may be similar to those of other metals which perturb protein function by binding to thiols,3,6 displacing native metal cofactors,99 or causing aggregation.7,100 Additional impacts on proteins may be related to KP1019’s effect on protein turnover. For example, our visualization of interactions between proteins with lowered abundance revealed a small cluster of proteins involved in the ubiquitin–proteasome system (Fig. 4A). Notably, deletion of RAD4 and RPT6 have previously been shown to increase accumulation of ubiquitinated proteins.101,102 Thus KP1019-dependent reductions in the levels of these proteins may contribute to the previously observed ability of the drug to result in global increases in ubiquitination.31

Regardless of the mechanism(s) by which KP1019 impacts protein folding and turnover, bioinformatics (YeastMine) analysis implicated the evolutionarily conserved transcription factor Hsf1 as a potential contributor to KP1019-dependent changes in protein levels. A reporter assay supported this conclusion (Fig. 8). In yeast and mammalian cells, Hsf1 is repressed by its interactions with Hsp70 chaperones.73,103 In S. cerevisiae, derepression can occur by misfolded proteins titrating Ssa1 (Hsp70) away from Hsf175 or by chemicals binding to or oxidizing Ssa1 thiols, thereby removing/displacing chaperones from Hsf1.37 Additional research will be required to determine which of these mechanisms, if any, applies to KP1019. The importance of drug-dependent perturbation of protein homeostasis relative to the drug’s genotoxicity and ROS-inducing capabilities also remains an open question, as HSF1 is an essential gene in yeast,104 which precludes analysis of deletion strains.

Beyond alterations to protein (re)folding and turnover, we observed significant increases in levels of proteins involved in translation (Fig. 3). Although many stresses have previously been shown to repress expression of genes involved in ribosomal structure and biogenesis,105 recent transcriptomic analyses of KP1019,12 other ruthenium complexes,106,107 and the metal salts AgNO₃, CdCl₂, HgCl₂, and ZnSO₄ showed induction of ribosomal structural and biogenesis genes. Moreover, a re-analysis of published transcriptomic data,29 using a 1.5-fold (vs. 3.5-fold) cut-off for gene induction also revealed an enrichment of ribosomal genes. Specifically, PANTHER identified “cytosolic translation” as the biological process GO term with the lowest FDR (0.0002). Notably, low levels (0.02%) of the DNA damaging agent methyl methane sulfonate (MMS) have also been shown to induce expression of ribosomal proteins in yeast.50 Although MMS is capable of damaging proteins,108–110 a major mode of its action is DNA damage, raising the possibility that KP1019's impacts on ribosomal proteins/genes may be related to its genotoxicity. The induction of ribosomal proteins following KP101912 and MMS50 treatment has been linked to the transcription factor Sfp1, which was implicated in our bioinformatics analysis (Fig. 5A). As a “generalist,” Sfp1 responds to diverse stresses, including oxidative and osmotic stress,111 both of which are caused by KP1019 (Fig. 6 and Singh et al.13). However, precisely how KP1019 impacts translation remains unclear. For example, it is not yet known whether KP1019 affects expression of ribosomal RNAs and/or whether the drug impacts the pool of mRNAs being translated (i.e. – the translatome).

Regardless, since elevated levels of translation machinery correlate with oxaliplatin sensitivity,86 we hypothesized that KP1019-dependent increases in abundance of ribosomal and ribosomal biogenesis proteins might sensitize yeast to oxaliplatin. In fact, that was the case (Fig. 9), which is consistent with models for synergy which predict that agents acting in similar processes may be more likely to have synergistic (as opposed to additive or antagonistic) effects.112 Given that both oxaliplatin and KP1019 impact translation (Fig. 3, ref. 32 and 86) and that “translation addicted” cancer cell lines are hypersensitive to oxaliplatin,86 future studies might explore whether these same cell lines are hypersensitive to KP1019. If so, KP1019’s effects on protein synthesis may explain its specificity for cancer cells, as many hematological and solid tumours have elevated levels of translation machinery.113–115 This specificity for translation-addicted cancer cells may, in turn, contribute to KP1019’s lack of dose-limiting toxicity.

Conclusions

Here we report the first global proteomic analysis of the effects of KP1019 on eukaryotic cells. Our findings reveal a multifaceted response that could be explained by the presence of both nuclear and non-nuclear targets. Although KP1019’s ability to damage DNA is well-established in both cancer and yeast cells,24,27,28 here we provide the first evidence supporting KP1019-dependent induction of oxidative stress in yeast. This finding further supports the utility of yeast as a model for studying KP1019, as this drug has been shown to cause oxidative stress in colorectal cancer cells.24 The observed increases in chaperones and translation machinery suggest that proteins serve as a significant cytosolic component of the drug. These findings may have long-term clinical implications, as translation has emerged as a promising target for chemotherapeutics.115,116 Moreover, the ability of KP1019 to sensitize yeast to oxaliplatin suggests that KP1019 may be a robust component of combination therapy regimens.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

This work was supported by a National Institutes of Health award to the UAB Comprehensive Cancer Center [grant number P30CA013148], by the UAB Institutional Core Funding Mechanism, by a National Science Foundation award to Birmingham-Southern College (1229016), and by the BSC rise³ stipend program. We would also like thank Stephen Elledge (Harvard University), Kevin Morano (University of Texas Health
Metallomics, 2020, 12, 876-890

References


This journal is © The Royal Society of Chemistry 2020

Metallomics, 2020, 12, 876–890 | 887


M. I. Webb and C. J. Walsby, EPR as a probe of the intracellular speciation of ruthenium(III) anticancer compounds, Metallomics, 2013, 5, 1624.


C. J. Gimeno, P. O. Ljungdahl, C. A. Styles and G. R. Fink, Unipolar cell divisions in the yeast S. cerevisiae lead to filamentous growth: Regulation by starvation and RAS, Cell, 1992, 68, 1077–1090.


A. Ianevski, L. He, T. Aittokallio and J. Tang, lacZ

L. Stultz, J. Mobley and P. Hanson, Budding yeast proteomic response to the anticancer ruthenium complex KP1019, Mendeley Data, 2020, DOI: 10.17632/vrsrwrjcy2.2.


81 V. M. Navadgi-Patil and P. M. Burgers, A tale of two tails: activation of DNA damage checkpoint kinase Mec1/ATR by the 9-1-1 clamp and by Dpb11/TopBP1, *DNA Repair*, 2009, 8, 996–1003.


